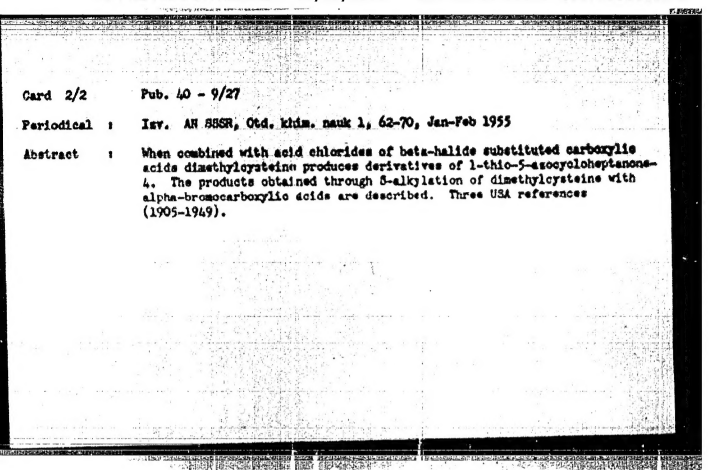
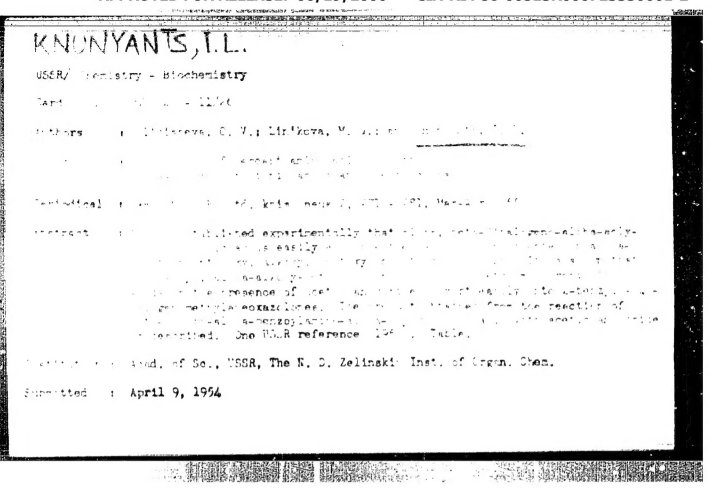
KNUNYANES, L.B. USER/ Chemistry - Conversions Card 1/2 Pub. A0 - 9/27 Eminyants, I. L., and Lin'kova, M. C. Authors Conversions of mercaptosmino acids. Part 2. Acylation and alkylation Title of dimethylcysteine Periodical Isv. AN SSSR, Otd. Khim. nauk 1, 62-70, Jan-Feb 1955 Abstract Experimental data are presented showing that disethylogateine anylates easily with acid anhydrides and acid chlorides resulting in the formation of only H-acyl derivatives. The aqueous-alkaline dimethylcysteine solutions alkylate easily, especially with halide substituted soids, forming only S-alkyl derivatives. Institution : Acad, of Sc., USER, The N. D. Zelinskiy Inst. of Org. Chem. Submitted : April 9, 1954



KNUNYANES, USSR/ Chemistry - Conversions Card 1/1 Pub. 40 - 10/27 Authors Emunyants, I. L.; Kil'disheva, O. V.; and Lin'kova, M. O. Conversions of mercaptosmino acids. Part 3. Acylation and alkylation Title of dimethylcysteins Periodical * Isy. AN SSSR. Otd. khim. nauk 1, 71-77, Jan-Feb 1955 Abstract The derivation of various M-scrylin derivatives of disethylogeteine containing Br, Cl and mathoxyl in the acyl radical is described. It is shown that the above mentioned derivatives cyclate as a result of the intramolecular attachment of the mercapto group of dimethylogeteine in place of the multiple bond of the crylic radical forming 1-this-5-ass-cyclo-heptanone -3. The results obtained from the reaction of dimethylogeteine with unsaturated acids and their derivatives, are explained. One USSR reference (1955). Acad. of So., USSR, The M. D. Zelinskiy Inst. of Org. Chem. Institution Submitted April 9, 1954

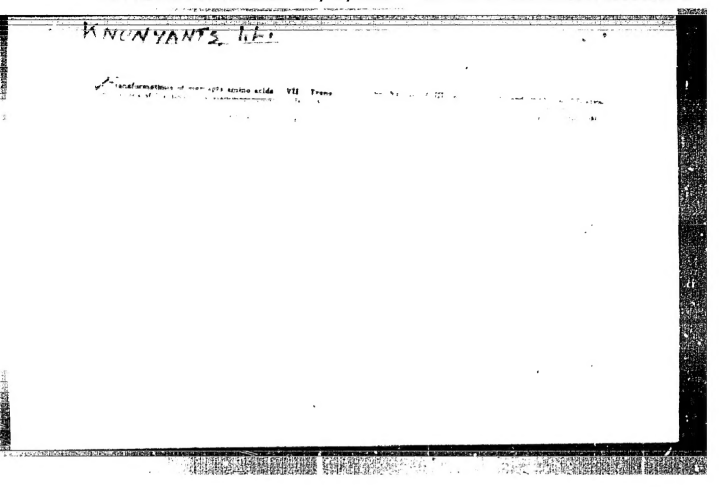
KNUNYANTS, I.L. USSR/ Chemistry - Biochemistry Pub. 40 - 10/26 Card 1/1 : Kil'disheva, O. V.; Restaykens, L. P.; and Knumyants, I. L. Au thore ! Conversions of mercapicamino soids. Part. 4. Alpha, beta-dihalogeno-alpha-T1 +.1.a acyl aminopropionic acids Periodical : Izv. AN SSSR. Otd. khim. nauk 2, 260 - 270, Mar-Apr 1955 I A study of the halogenation reaction of alpha-acylaminoacrylic acids showed that they combine easily with Cl and Br forming sufficient quantities of Abstract s, reta-divalogence-algha-acylamic or the notice. The most feverable . Will or Br solutions in dry chloroform or carren fetoschoride redia. withat reta-dibalogeno-alpha-adylamingtrorients acts see fount incolucie in oviss and 2 English (1930-1954). Tobler. Institution : Acad. of Sc., "SSR, The N. D. Zelinskip Inst. of from. Der. Submitted : April 9, 1954

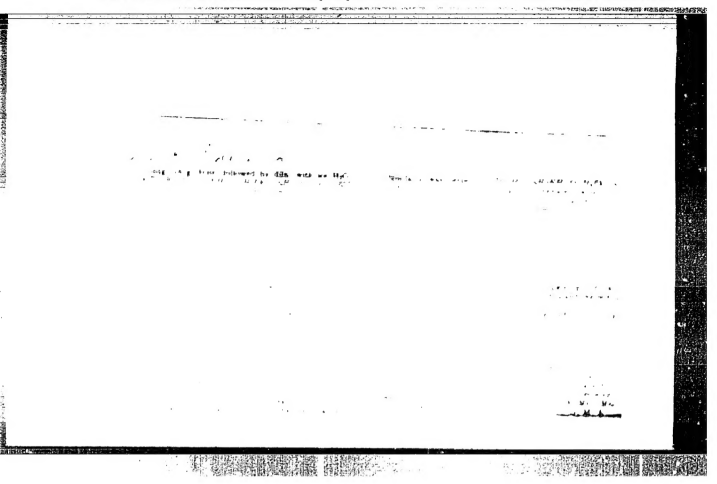


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CIA-RDP86-00513R000723330001-2

VANTSILLI USSR/ Charistry - Biochemistry Card 11 Pub. 40 - 12/26 1 Ell'disheva, O. V.; Lin'kova, M. G.: and Enunyants, I. L. tu thors I contel to of moroaytoamine acits. Far F. Setonia comments almomany lumino on ordin actins and their derivatives Periodical : Isv. AN SSSR. Otd. khim. nauk 2, 262 - 288, Mar-Arr 1955 instract : It is shown that thermal cleavage of hydroson callie from all ha, beta-dihaloyes -all a-acylaminoprogionic acti results to the formation of bet -balternore-arglamineacrylic acid. The room of motion from the relation of the mylic soid with acetic augminute is to a sed. The derivation of the Siret training of meatureted tetames accounties form minit set a tra terms, amountaide of almha-combongemino-pain-transfer in the first the reaction of beta-bromo-alpha-a rhocorzylog or eartier's is discreed. Six ref renners a con, of ic., 'S A, The N. D. Dellis, 'S and a Submitted : Abril 9, 1954

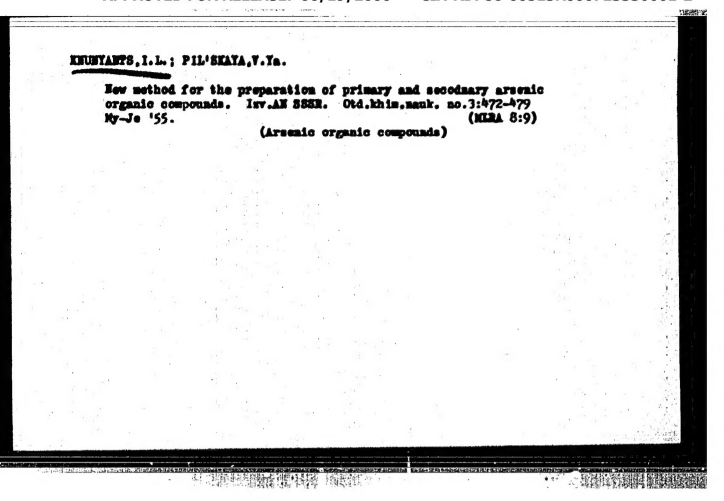




TRUSTARTS, I.L.: EHCHMA, V.V.

Conversions of mercapte amine soids. Report no.8. Alkylation and acylation of cysteins and dinethylogeteins with derivatives of cy-acylamino-β-halogenopropicaic soids. Inv.AB SSER. Otd. hhim.namk no.3:462-471 My-Je '55. (MERA 8:9)

1. Institut organicheakoy khimit in. N.D.Zelinskogo Akademit namk SSER. (Gysteins) (Propionic soid)



LIB'ROVA, M.G.; RIL'DIERWA, O.V.; REUBTARTS, I.L.

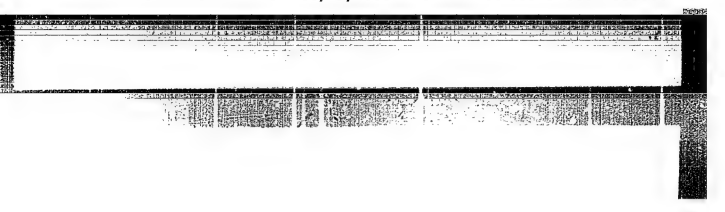
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REUFTANTS, I.L.; RIL'DISHEVA, O.V.; PERTOVA, To. To.

Conversion of mercaptesmine acids. Report no.10.New method for the synthesis of polypoptides. Isv.AH SSER.Otd.khim.mank no.4: 696-704 J1-Ag 155. (NIBA 9:1)

l. Institut erganicheskey khimii imeni W.D. Zelinskege Akademii nauk SSER.

(Poptides) (Amine soids)



KNUNYANTS, I.L.

AID P - 3159

Subject

g USSE/Chemistry

Card 1/1 Pub. 119 - 1/7

Authors

! Knunyants, I. L. and Ye. Ta. Pervova (Moscow)

Title

1 Progress in establishing the structure of proteins and their

synthesis

Periodical : Usp. khim., 246, 641-672, 1955

Abstract

1 Methods of protecting the amino group during acylation as well as for the prevention of condensation are reviewed. Synthesis of numerous polypeptides is described. Several protecting agents are mentioned. The review is based on non-Russian literature exclucively.

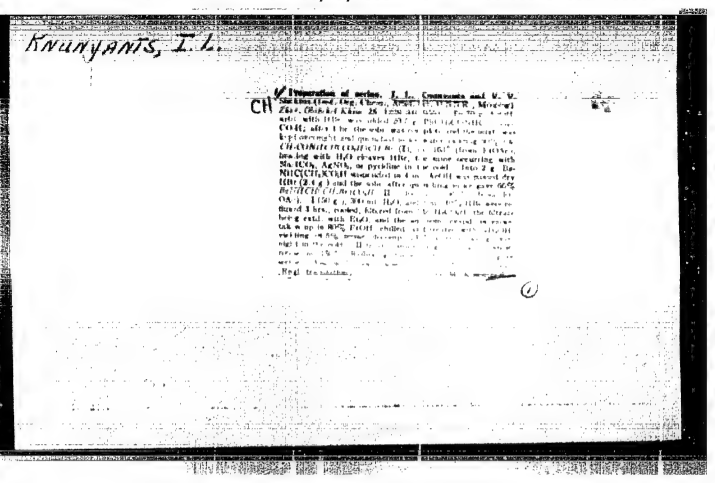
Institution: None

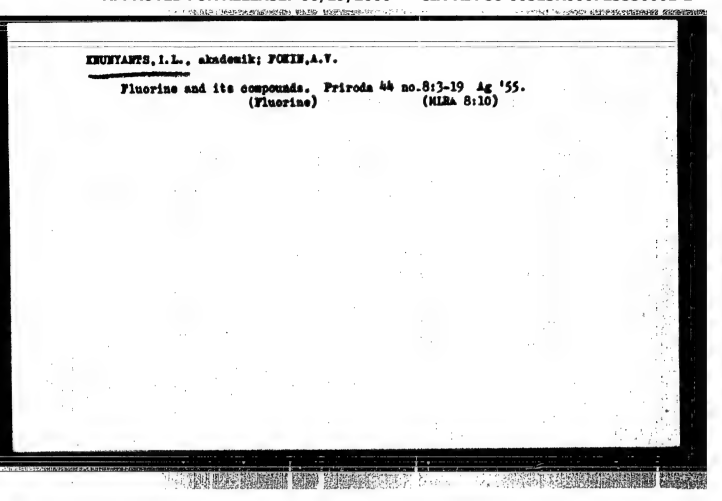
Submitted : No date

SHOKINA, V.V.; KMUSTANES, I.L.

Aldehydic acid halydes. Shur.ob.khim. 25 no.4:758-760 Ap '55. (HIRA 8:7)

l. Institut organicheskoy khimii Akademii Hauk SSSR. (Aldehydic acids) (Halides)





SINORS, J.H., editor; Eminyants Libra akademik, redaktor; VARSHAVSKIY, Ya.M., kandidat khimicheskikh nauk, redaktor; MARSHAVSKIY, V.A., redaktor; GRIBOYA, N.P., tekhnicheskiy redaktor

[Fluorine chemistry. Translated from the English] Ftor i ego scedimentia. Perevod s anglitskogo. Pod red. I.L.Enuniantsa i IA.M. Varshavskogo. Moskva, Isd-vo incetrannoi lit-ry, Vol.2. 1956, 495 p.

(Fluorine)

可图式作用 医电阻性视器 医氯酚烷酸 医红色性原体点点

The parties of the second seco

RODIOMOV, V.N., akademik, redaktor [deceased]; KAZARSKIY, \$.A., akademik, redaktor; EKMYAKUYS, I.J., akademik, redaktor; SEMYAKUY, N.N., redaktor; HAL'HIKOV, N.N., professor, redaktor; TAYYS, S.Z., redaktor; SHMASTINA, Ye.V., redaktor; KORMYRYA, V.I., tekhnicheskiy redaktor

[Resctions and methods of analysis of organic compounds] Reaktsii i metody issledovaniia organicheskikh seedinenii. Noskva, Gos. nanehnotekha, isd-ve khim, lit-ry. Vol.4, 1956, 319 p. (MEMA 917)

1. Ohlen-korrespondent AN SEER (for Shenyakin)
(Chemical reactions) (Isomers and isomerization)

Card 1/1

- 115 -

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R00072333000

EMUMYANTS, 1.1.; PERVOVA, Te.Ta.; STURMENVA,V.V.

Reactions of perfluere elefins. Part 5. Reactions for the conjugate addition of halides, Isv.AN 888R Otd.khim.mank no.7:843-849 Jl 156.

1. Institut elementeergamicheskikh seyedinemiy Akademii nauk 285R. (Olefine) (Malides)

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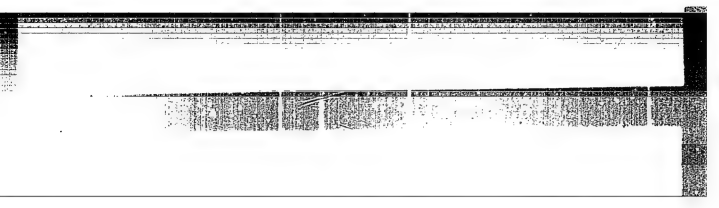
B-2

USSR/Organic Chemistry, Synthetic Organic Chemistry.

Abe Jours Ref Zair-Khilelya No 6, 1997, 19069

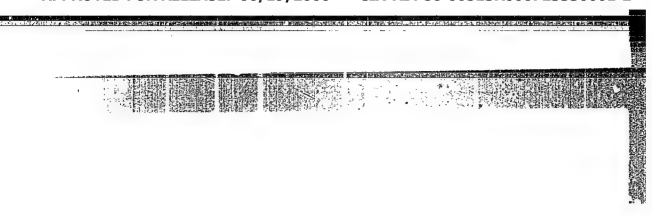
dimethylacrylic acid, are treated with a solution of 7.1 g. Clo in 71 cc CCl. I is obtained, yield 93%, melting point 10% (decomp.). 7.5 g. I are treated with 30 cc of water, after 2 hours the precipitate is filtered off, yield II 73%, m. p. 70-75° (dec.). A mixture of 5.8 g. I, 50 sc(CHqUO)gO, and 25cc CCli is heated in a vacuum at 70-80° IV is obtained, yield 65%, m.p. 95° (from absolute sp). Analogically, at the treatment of 2.4 g. II with 25cc (CHgCO)gO in CCli, IV is obtained, yield 80%. At the action of III cm I in C6H6 the yield of IV is 20%. From the filtrate after treatment with bicarbonate II, m.p. 155° precipitates. 0.01 mole I heated for 2 hours with 0.01 mole PCls in 20 cc abs. C6H6 is produced 2-phenyl-1-(2-chloro-isopropyl)-commolone (VI), yield 72%, m.p. 60-64° (decomp., from petr. ether).0.01 mole I is treated with a solution

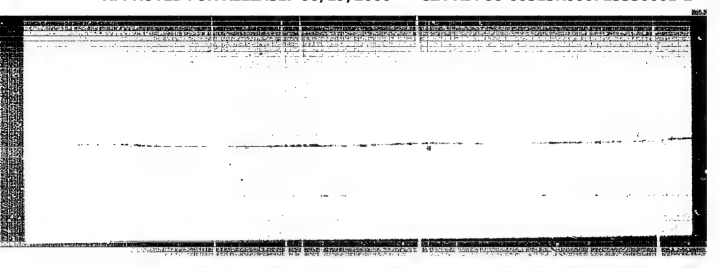
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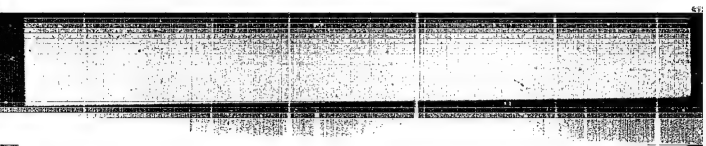












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USSR/Organic Chemistry - Theoretical and General Questions on Organic Chemistry

E-1

Abs Jour

: Referat Zhur - Khimiya, No 2, 1957, 4210

Author

: Knunvants, I.L., Dyatkin, B.L., Gambaryan, N.P.

Title

: OnReady and Widespread Formation of Four-Membered Ring

Orig Pub

: Uspekhi khimii, 1956, 25, No 7, 785-844

Abstract

: A synopsis of numerous literature data concerning the preparation and the stability of four-membered cyclic compounds. It is shown that ready formation and stability of four-membered rings are substantially affectod by the presence of substituents and their nature. In a number of cases formation of four-membered rings takes place in preference to that of the five- and six-membered cyclic systems,

Bibliography 414 references.

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001

KNUNGANIS, 2, L

In memory of A.A. Strepikheev; obityary, Thur.ob.khim.26 no.11:3224-3226 H 156. (Strepikheev, Aleksandr Aleksandrovich, 1912-1955) (MIRA 10:1)

USSR/Organic Chemistry. Synthetic Organic Chemistry:

Abs Jours Mererat Zimr-Khimiya, No 4, 1958, 11392.

Author : Minyante Citt and Pokin, A. V. Inst Academy of Sciences USER

Title : The Hitration of Perflucroolefins by Mitrogen Dioxide

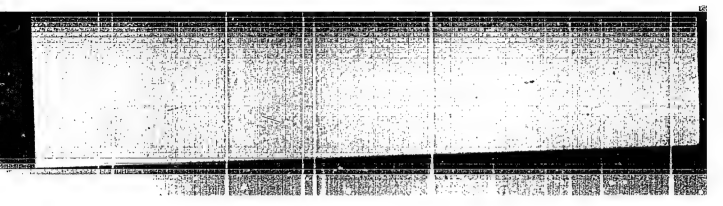
Orig Pub: Doklady Akad Heuk MISR, 111, No 5, 1035-1038 (1956) \$4.40 10.50 Pan 海绵湖湖湖湖湖湖流水流 (1.20 ex 10 sector 10 se

Abstract: The reaction of Egogavith perfluctoolefins proceeds by a free redical mechanism and leads to the forma-tion of diniaroperfluctoolkanes and & mitroperflucto-alkylnitrites; the overall yield is 90%. The reactivity of the perfluorcolefins decreases from left to right in the following series OFg-CF2 (I) > CF3CF-CF2 (II) > In occlusion of the control of the contr

: 1/3 Card

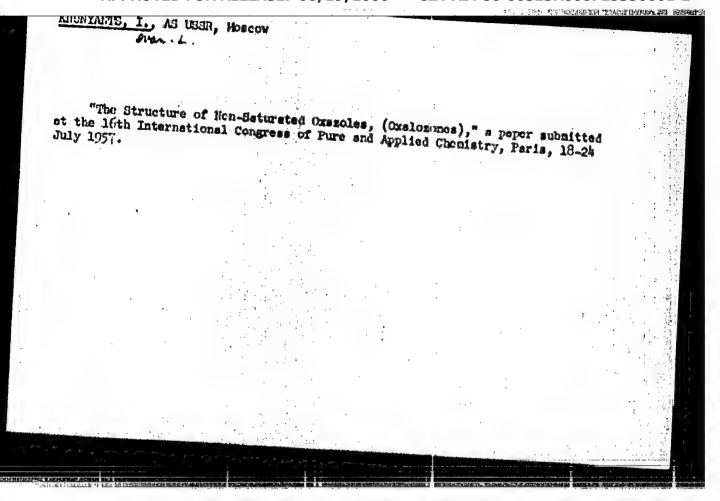
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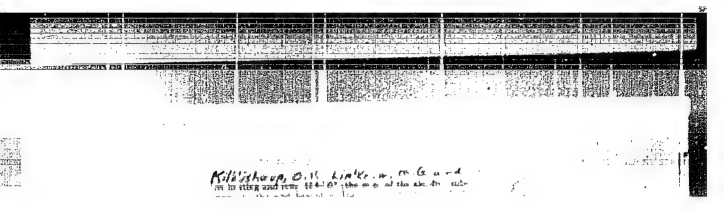


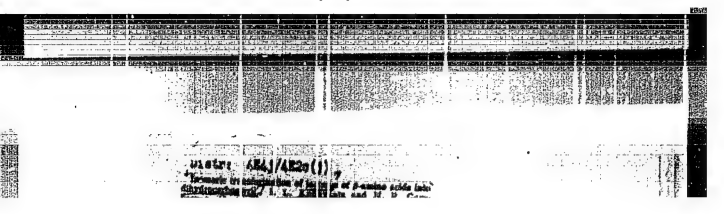
TERENT'INV, A.P.; YANOVSKAYA, L.A.; RUENARE, Ye.O., redaktor;
RODIONOV, V.M., akademik, redaktor [deaceased]; KARANSHI, B.A.,
akademik, redaktor; Kunyappa L.L., akademik, redaktor;
SHEMYAKIN, M.M., redaktor; Mil. Birov, M.T., prof. redaktor;
LURYE, M.S., teknicheskiy redaktor.

[Polarographic analysis in organic chemistry] Poliarograficheskii
method v organicheski khimii. Moskva, Gon. nauchno - teknic isday
vo khim. lit-ry, 1957. 368 p. (Reaktsii i metody iseledovaniia
organicheskikh soedimenii, vol.5) (MIRA 10:10)

1. Chlen-korrespondent am SSSR (for Shemyahin).

(Polarography) (Chemistry, Organic)





A-1

KNUY ANTS, L.K.

USSR / General Topics. Methodology, History, Scientific Institutions and Conferences, Instruction, Bibliography and Scientific Documentation.

Abs Jour : Ref Zhur - Khimiya, No 5, 1958, No 13h10

Author & L.L. Knuyants

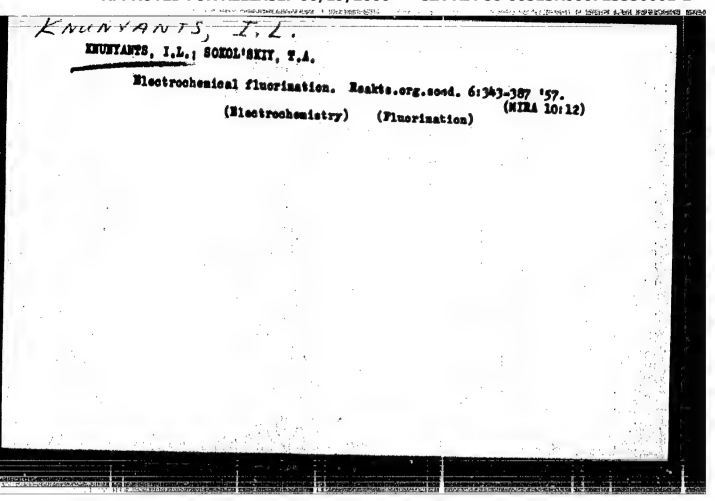
Inst : Not given

Title : Basic Development Trends of Soviet Science of Chemistry

Orig Pub : Khim. nauka 1 prom-st', 1957, 2, No 5, 538 - 569

Abstract : To the 40th Anniversary of the Great October socialist revolution. A review of basic divisions of chemistry in 40 years.

Card : 1/1



KIL'DISHEVA, O.V.; LIN'KOVA, M.G.; EMUSYANTS, I.L.

Structure and properties of oxasolones, lav. AN ESER Otd. khis. mark no.6:719-729 Je '57. (MIRA 10:211)

1. Institut elementoorganicheskikh soyedineniy AN SEGR. (Oxasolone)

AUTHORS:

Knunyants, I.L., Fokin, A.V.

62-12-3/20

TITLE

Mitration of Fluorine Olefines by Mitrogen Dioxyde (Mitrovaniye) fternlefinov dvuckis'ju azota). Lecture Delivered at the Lecting of the Department of Chemical Sciences AN USER on Cotober 30, 19 (Doklad na sessii Otdeleniya khimicheskikh nauk Akademii nauk SSSR 30 oktyabrya 1957 g).

Investiga AN SSSR Otdelenie Khimicheskikh Neuk, 1957, Hr 12,

pp. 1439-1451 (USSR)

ABSTRACT:

PERIODICAL

The reactions of the nitration of saturated and unsaturated hydrocarbons, which have already been thoroughly investigated, were practically not investigated at all in fluorine-organic compounds. The majority of the reaction of fluorine olefines with "mucleophylic" reagents has ionic character. In contrast to the smooth interaction of perfluorine-olefines with nucleophylic reagents, their interaction with electrophylic substances takes place under much more difficult conditions. According to Koffman (1949) and Khastsel'din (1953) only dinitroalkanes are formed by the interaction of tetrafluorine ethylene and chlorine trifluorineethylene with nitrogen dioxide. Reactions with other fluorine clefines were not investigate.

Card 1/3

ed. Among other things, the authors found that these reactions

Mitration of Fluorine Olefines by Mitrogen Dioxyde. Lecture Delivered at the Meeting of the Department of Chemical Sciences AN USSR on October 30, 1957

62-12-3/20

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depend mainly on the structure of the characteristic features of fluorine olefine (and lead to the formation of new and interesting substances). It was further shown that tetrafluorine-ethylene reacts explosively with nitrogen dioxide. It was possible to extend the method of nitration by means of nitrogen-dioxide also to other perfluorine olefines (see tables). In the case of none of the methods investigated were compounds able to form. It was shown that the destruction of 3-nitroperfluorine-ethyl and 3-nitroperfluorpro-pylnitrites begins only at a temperature of more than 2500. The investigation of the nitration of chlorine fluorine elefines made it possible to determine a certain characteristic feature of this reaution (see formulae on page 1446.) The investigation of the nitration reactions of fluorine olefines and not substituted olefines with nitrogen-dioxide made it apparent that there is a similarity of the chemical character of these reactions (see table 2). The results of this investigation further showed that the stability of intermediate radicals as well as the polarity of fluorine olefines and that of the radical-like particle NO2 are an important factor

Card 2/3

Nitration of Pluorine Olefines by Mitrogen Dioxyde. Lecture Delivered at the Meeting of the Department of Chemical Sciences AN USSR on October 30, 1957 62-12-3/20

of orientation of the reacting components. Conceptions concerning the polarity of radicals, which were first published by Ucters (waters?) and were further developed by Karash, Veys, Dolgopolov and others, deserve attention. There are 2 tables and 18 references, 10 of which are Slavic.

SUBMITTED:

October 9, 1957

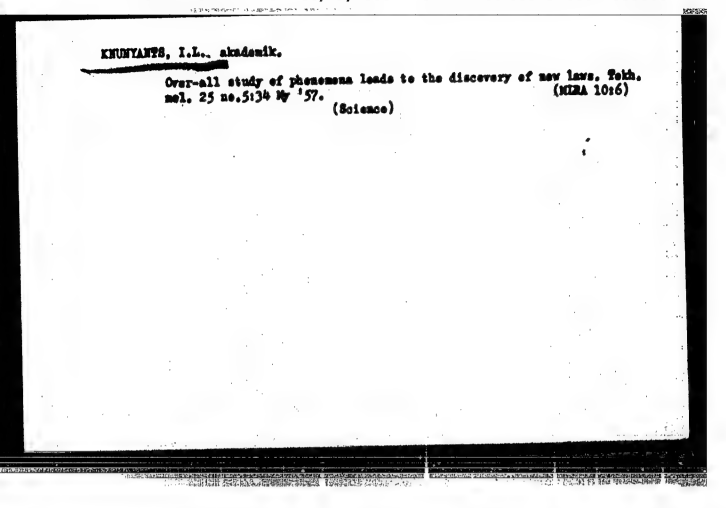
AVAILABLE:

Library of Congress

Card 3/3

1. Chemical engineering-Conference 2. Hydrocarbon-Reactions

3. Fluorine-Organic compounds 4. Fluorine clefines



AUTHOR TITLE

ABSTRACT

PA - 2914 EMUNYANTS, I.L., Member of the Academy, VYAZANKIN, N.S. Reduction Dimerisation of Derivates of a, 8-Unsaturated Acids. (Vosstanovitelnaya dimerisatsiya proisvodnykh α,β-nenasyshohennykh

kislot -Russian) PERIODICAL

Doklady Akademii Hauk 8558, 1957, Vol 113, Mr 1, pp 112-115, (U.B. 8.8.)

Received 6/1957

As already dated, dimitryle of adipine acid forms the main product of the electrochemical reduction of acrylomitryle as it was already mentioned. This work was carried out in order to investigate the dependence of hydrodimerisation phenomena on the etructure of the compounds to be reduced and on the nature of the analgams. The investigation showed that one of the factors which influence the formation of hydrodinerigates is the character of the conjugated system of the substance to be reduced. This is by no means unexpected if it is assumed that the formation of the hydrodimerigate is preceded by molecule reduction withits subsequent dimerization. It is known that the non-conjugated ethylene-bindings to analgam are not reduced. It was found (in the case of stirol and poyvinacetate) that the conjugations of the ethylene binding with the bernol kerml or with one unseparated electron-couple of the coygen atom are not sufficient even for the reduction of a double-binding to potassium or lithium analgam. It is rather difficult to reduce amides and diethyl smides of the same soid, they give no hydrodimerisates, Diethylamide and diphenyleamide of cinnamon acid produce, besides the normal products of the reduction, also hydrodi-

Card 1/2

· KNUNYANTS, I.L.

20-2-24/60

Dyatkin, B. L., German, L. S., Knunyants, I. L., Member

AUTHORS: of the Academy

Anionotropic Rearrangement of Substituted Perfluoropropenes TITLE:

(Anionotrophaya peregruppirovka sameshchennykh perftorpro-

penov)

PERIODICAL:

Doklady Akademii Hauk SSER, 1957, Vol. 114, Hr 2, pp. 320-322

(USSR)

ABSTRACT:

As was shown by the authors of the paper under review in an earlier scientific publication, the reactions of affiliation and of vinylic substitution are in competition with each other if we have the case of an interaction of perfluoropropylene and perfuorisobutylene, on the one hand, with alcohols and amines, on the other hand. No allylic substitution takes place. This demonstrates that in the molecules of these fluorolefines the effects of conjugation of the double bond with the Q-F-bond in the CF3-group are weak. From this point of view, the reactions of the chlorofluoropropenes and chlorofluorobutenes command great interest, particularly the reac-

Card 1/3

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APPROVED FOR RELEASE: 06/19/2000

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20-2-24/60

Anionotropic Rearrangement of Substituted Perfluoropropenes

tions of perfluorallylchloride CF2=CF-CF2Cl. There also exists information according to which influence of nucleophile reagents on perfluorallylchloride leads to a substitution of chloring by a corresponding anion. It has to be assumed that this is the result of the conjugation of the bond C-Cl with the double bond. The authors of the present paper investigated the reactions of perfluorallylchloride with sodium methylate and diethylamine. The interaction with the sodium methylate leads to the perfluorally lathylether. This reaction represents a new solution for arriving at the derivatives of the perfluoracrylic acid. The ether is polymerized even at a lower temperature. The reaction of perfluorallylchloride with dethylamine has a light course. The perfluorallyldiethylamine produced as result of this reaction rearranges itself, still during the reaction, into perfluorpropenyldiethylamine, Hydrolysis of the latter leads to diethylamide of the oc-hydroperfluorpropionic acid. Bromination of the perfluorpropenyldiethylamine with a subsequent hydrolysis results in diethylanide of the Ct-bromoperfluorpropionic acid. The above isomerisation represents an allylic rearrangement and probably is caused by the tendency towards formation of a stabler system, and this owing to the conjugation of a double bond

Card 2/3

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-

Anionotropic Rearrangement of Substituted Perfluoropropenes

with an unseparated electron pair of the substituent in the allylic position. The velocity of the rearrangement depends of the degree of mobility of the electron pair. Different compounds are arranged in a series in accordance with the criterion of stability. The experimental part of the paper under review contains the production methods together with the constants and yields of the substances investigated. There are 3 references, 1 of which is Soviet.

SUBMITTED:

January 18, 1957

AVAILABLE:

Library of Congress

KNUNYANTS, I-L.

5(3)

PHASE I BOOK EXPLOITATION

SOV/1589

Akademiya nauk SSSR.

Khimiya bol'shikh molekul; sbornik statey (Chemistry of Large Molecules; Collection of Articles) Moscow, Izd-vo AM SSSR, 1958. 299 p. (Series: Akademiya nauk SSSR. Mauchno-populyarnaya seriya) 30,000 copies printed.

Compiler: G.V. Sklovskiy; Resp. Ed.: A.V. Topchiyev, Academician; Ed. of Publishing House: V.A. Boyarskiy; Tech. Ed.: I.N. Guseva.

PURPOSE: This book is intended for a wide circle of readers including those who have had no training in chemistry. It can also serve as amanual for propagandists, teachers, and journalists.

Card 1/8

Chemistry of Large Molecules (Cont.)

307/1589

COVERAGE: This collection of articles reflects the trend for the future development of the Soviet chemical industry as indicated by the May plenary session of the Central Committee of the Communist Party within the framework of the new Seven Year Plan: These articles were published in newspapers and journals. The authors, scientists and industry workers, developed the theme of accelerated development of the chemical industries, and sciences, with stress on the manufacture of synthetic fibers, plastics, and other materials. Some of the articles were abridged, revised, or enlarged. The articles were selected so as to give an adequate survey of the chemistry and technology of high-molecular-weight compounds and their use in industry, agriculture, and in the manufacture of consumers' goods. Mantioned are raw materials for the production of polymers. This book belongs to the popular-science series of the Academy of Sciences. Similar volumes are intended for future publication. No references are given.

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RODIOHOV, Vladimir Mikhaylovich, akademik [deceased]; SVORTKHA, V.I.,
sostavitel'; KISMENA, V.V., sostavitel'; FEOROVA, A.M.,
[translator]; KUNTANTS, L.L., akademik, ctv.red.; SERNTAKH, N.M.;
akademik, ctv.red.; SHVEROV, Tu.B., red.ind.; FOLDMOVA, 2.P.,
tekhn.red.

[Selected works] Isbrannye trudy. Moskva, Ind-vo Akad. mank SSSE,
1958. 792 p.
(Chemistry, Organic)

(Chemistry, Organic)

HRUFTH 78, [.]. PHYOVA, To. Za.; LIH'ENVA, N.G.; KIL'DISHWA, O.V.

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STRELLY, R.W.; YATSENKO, R.D.; KREYANTE.

Reaction of perfluorovinyl magnetium iodide with carbonyl compounds.

(NIRA 11:10

(Vinyl compounds) (Carbonyl compounds) (MIRA 11:10)

> reacts with a symmetric fluoroletins, only one of the isomers forms which contains the difluormethylene group in the

There is 1 table and 4 references, 1 of which is Soviet,

APPROVED FOR BELLASE; 06,019/2000

CIA-RDP86-00513R000723330001-2

Card 1/1

THE STATE OF THE CASE AND DESCRIPTION OF THE PERSONS

SOV/63-3-6-30/43

AUTHORS: Knunyants, I.L., Dyatkin, B.L., Gorman, L.S.

TITLE: Reactions of Perfluoroscrylonitril (Reaktsii perftorakrilo-

PERIODICAL: Khimicheskaya nauka i promyshlennost', 1958, Vol III, Hr 6,

pp 020-029 (USSR)

ABSTRACT: It has been shown that pure perfluoroscryionitril easily re-

acts with methanol and ethanol producing β -alkoxy- α -hydroperfluoropropionitiils. It reacts also with piperidim anianilin in an ether solution producing amine fluorohydrate. There is I table and 2 non-Soviet references.

ASSOCIATION: Institut elementoorganichookikh soyedineniy Akademii nauk SSSR

(Institute of Elemental - Organic Compounds of the USSR Academy of

Sciences)

SUBMITTED: July 10, 1958

Card 1/1

Card 1/2

AUTHORS. 62-58-3-6/30 Sterlin, R. N. , Dyaskin, B. L. Pinkina, L. R. , TITLE: Reactions of Fluorolefine: (Reakteil ftorolefinor) Communication 7. Addition Compounds of Acid Chlorides to Vinylidene Fluoride and Trifluoroethylene (Soobshcheniye 7. Prisoyedineniye khlorangidridov kislot k floristomu vinilidenu i triftoretilenu) PERIODICAL: Izvestiya Akademii Nauk SSSR, Otdeleniye Khimicheekikh Nauk, 1958, Nr 3, pp. 296 - 299 (USSR) Street Property Lines ABSTRACT: The addition of alkyl halides discovered by Kondakov was later developed by others. In the present paper the authors show that such fluorolefine as vinylidene fluoride and trifluorethylene (in the presence of nonaqueous AlOl,) possess the capability of combining with carboxyl chlorides and thereby forming the corresponding fluoro-substituted ketones. Vinylidene fluoride very readily combines with the acid chlorides

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-2"

of butyrig acid and propionic acid at a temperature of

-5 to -10°C in the presence of equivalent quanta of AlCl in

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Reactions of Fluorolefins. Economication: 7. Addition Compounds of Acid Chlorides to Vinylidene Fluoride and Trifluoroethylene

pure chloroform, where alkyl-2-chloro-2,2-difluorosthyl-ketones with yields of 44,48: and 33 % form:

At the same time substances form which correspond to the products of the partial or complete substitution of fluorine in chlorine and the products of further condensation. The authors obtained: methyl-2-chloro-2,2-difluoroethylketone and methyl-2-chloro-1,2,2-trifluoroethylketone. There are 8 references, 2 of which are 80viet.

SUBMITTED:

November 3, 1956

Card 2/2

. MUNYANTS, IL.

62-58-4-6/32

AUTHORS:

Knunyants, I. L., Sterlin, R. N., Bogachev, V. Ye.

TITLE:

Reaction of Pluorolefines (Reaktsiy ftorolefinov) Communication 2. The Synthesis of 2-Iodoperfluorpropylene and Some of its Properties (Soobshcheniye 2. Polucheniye i nekotoryye svoystva 2-yodperftorpropilena)

PERIODICAL:

Izveitiya Akademii Nauk SSSR, Otdeleniye Khinicheskikh Hauk, 1958, Nr 4, pp. 425-427 (USSR)

ABSTRACT:

The most simple way of producing the second member of the perfluorowinyl-iodide series would be the dehydrohalogenation of 1-chloro-2-iodo-2-hydroperfluoropropane. This can be reached by interaction of 2-hydroperfluoro propylene with iodine chloride -. From papers dealing with the compounds of alcohols with fluorolefines the conclusion can be drawn that with the increase of the polarity of the olefines the alcohols more easily combine with these fluorolefines. There are, however, no concrete data on the number

Card 1/2

of references. The authors describe in this paper

62-58-4-6/32

TO THE PART SERVICE STREET, OF THE SERVICE ST

Reaction of Pluorolefines. Communication 2. The Synthesis of 2-Iodoperfluoropropylene and Some of its Properties

the synthesization of perfluoroisopropenyl iodide by combination of ethyl alcohol with perfluoroisopropenyl iodide. The earlier not described 1-methoxy-2-10do-2-hydroperfluoropropane was produced. By saronafication of this ester methyl esters of the &-iodo-&-hydroperfluoropropionic acid was produced. By dehalogenating this substance the methyl ester of 2-diffunroalkylic acid was produced. There are 7 referen-

ASSOCIATION: Institut elementoorganichenkikh soyedineniy Akademii nauk SSSR (Institute for Elemental-organic Jonyoundo, AS USSR)

SUBMITTED:

November 20, 1956

AVAILABLE:

Library of Congress

Card 2/2

1. Fluorelefines--Reaction

等。這門 強調 實際 特別

2. Iedoperfluorepropylene - Preperties

AUTHORS:

Knunyants, I. L., Dyatkin, B. L.

62-58-5-25/27

L. & PURITY PERMITTAL VIBRICATION PORTER SCENE

TITLE:

Interaction of Some Fluorine Olefine With Phenols (Vsaimodeystviye nekotorykh ftorolefinov s fenolami)

PERIODICAL:

Isvestiya Akademii Hauk SSSR, Otdeleniye Khimicheskikh Hauk 1958, Hr 5, pp. 648-650 (USSR)

ABSTRACT:

The special character of the binary bond of perfluoroisc-butylene and perfluoracrylonitryle consists among other of the fact that these compounds are easily subjected to an interaction with nucleophile reagents. Alcohols associate in this way without alkaline catalysts and without heating. In the case of perfluoroisobutylene, a substitution of the fluorine-vinyl atom into the alkoxy-group takes place. It was therefore of interest to investigate the behavior of these fluorolefines with respect to phenols. In the present report the authors describe the carried out non-catalytic compound of phenol and hydroquinone with fluoroisobutylene and perfluorakrylonitryle. Concluding, the description of the obtaining of phenylper-fluorallylester by the interaction of perfluorallylehloride with potassium phenolate is given. There are 1 table and 5 references, 2 of which are Soviet.

Card 1/2

Interaction of Some Fluorine Olefine With Phenols

62-58-5-25/27

ASSOCIATION:

Institut elementoorganicheskakh soyedineniy Akademii nauk

SSSR (Institute for Michella Laborganie Compounds AS USSR)

SUBMITTED:

January 4, 1956

1. Ethylenes-Chemical reactions 2. Phenols-Chemical reactions

Card 2/2

CIA-RDP86-00513R000723330001-2" APPROVED FOR RELEASE: 06/19/2000

AUTHORS:

Knunyante, I. L., Mysov, Ye. I.,

807/62-58-7-24/26

TITLE:

The Catalytic Hydration of the q-Olefines (Kataliticheskoye gidrirovaniye q-olefinov)

PERIODICAL:

Isvestiya Akademii nauk SSSR, Otdeleniye khimicheskikh nauk, 1958, Nr 7, pp. 906 - 907 (USSR)

ABSTRACT:

The investigation of the catalytic hydration of the e-clefines besides its practical importance is also interesting because it is directly connected with important problems concerning the theory of heterogeneous catalysis. The rate of hydration depends on the state of the m-bond of the clefines. It increases with the decrease of the electron density of the bond, if the removal of the electrons from the catalyst lattice by the clefine molecule is the primary phenomenon in this process. The authors of the present paper showed that e-sthylene, e-propylene, e-isobutylene, and other e-clefines may be easily hydrated with molecular hydrogen on a palladium and nickel catalyst. The enclosed table gives the formulae of the initial clefine, the name of the catalyst, the hydration temperatures, the hydration

Card 1/2

The Catalytic Hydration of the q-Olefines

SOY/62-58-7-24/26

products etc. Based on the observations made it may be assumed with great probability that the property of easy hydration of the olefine series increases from a-athylene to e-isobutylene. There are 1 table and ? non-Soviet references.

ASSOCIATION:

Institute of Elemental-Compounds, AS USSE)

SUBMITTED:

May 12, 1958

Card 2/2

在任存特殊的的指接**用的第三人称单数形式** KNUNYANTS, I.L. AUTHOR: None Given SOY/30-58-8-9/43 TITLE: At the Plenary Meetings of Departments (Na obshchikh sobraniyakh PERIODICAL: Vestnik Akademii nauk SSSR, 1958, Nr 8, pp. 57-68 (USSR) ABSTRACT: These plenary meetings were held on June 16-17 with the purpose of nominating candidates for the AS USSR. Scientific lectures Department of Physical and Mathematical Scir.ces: The Corresponding Member, AS, USSR, Ya. B. Zel'dovich spoke about the catalysis of nuclear reaction by mesons and the resulting phenomena. This hypothesis of a possibility of such a catalysis was first mentioned in 1954 by A. D. Sakharov and Ya. B. Department of Chemical Sciences: V. A. Kargin, Member, Academy of Sciences, USSR, spoke about the tasks and aims of the work of the Council of Scientists (uchenyy sovet) on polymeric compounds; the council consists of 6 sections: for the synthesis of monomers; for the synthesis and kinetics of reactions; for the recovery of polymeric compounds; for materials for air-Card 1/5 craft construction and other special polymeric materials; for

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At the Plenary Meetings of Departments

50V/30-58-8-9/43

chemical fibres; for the use and processing of polymeric materials. The council counts more than 100 persons. It comprises nearly all the leading scientists and experts of the respective branches of the chemical industry and of the universities. A. V. Topchiyev, Member, Academy of Sciences, USSR, reported on the working results of the commission for the elaboration of long-reads plans for scientific research to be conducted in the institutes of the AS, USSR, in the fields of the production and the use of high-molecular compounds. A lecture on cancerolytic peptides was held by I. L. Knunyants, Member, Academy of Sciences, USSR. This work which he carried out together with M. G. Golubeva and O. V. Kil'disheva is dedicated to the principal problems of cancer etiology. As suggested by the plenary meeting a special conference on that issue should be held under participation of a wide circle of physicians, biologists and chemists. Department of Geological and Geographical Sciences: N. S. Shatskiy, Member, Academy of Sciences, USSE, spoke about movements of the earth crust and their origin, and the Corresponding Member. Academy of Sciences, USSR, Y. Y.

Card 2/5

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At the Plenary Meetings of Departments

807/30-58-8-9/43

Belousov reported on some results and prospects of tectonicphysical investigations. Department of Biological Sciences: The plenary meeting was held at the new station (equipped with an air-conditioning plant) of the Institute of Plant Physiology imeni K. A. Timiryasev. A. L. Kursanov, Member, Academy of Sciences, USSR, and I. I. Tumanov, Corresponding Member, Academy of Sciences, USSR, spoke about their investigations in plant physiology at this station, AS USSR, equipped with an air-conditioning plant. V. M. Sukachev, Member, Academy of Sciences, USSE, presented new data concerning the experimental investigation of plant interrelations. Sukachev is the Bead of the Laboratoriya lesnoy geobotaniki Instituta lesa Akademii nauk SSSR (Laboratory for Forest-Geobotany at the Forestry Institute, AS USSR) at which this work was carried out. Ye. M. Mishustin, Member, Academy of Sciences, USSE, spoke about soil microorganisms. Department of Engineering Sciences: G. I. Petrov, Corresponding Member, Academy of Sciences, USSR, talked on motions in real gas with velocities exceeding by far the velocity of sound. V. S. Pugachev, Doctor of Technical Sciences, spoke about

Card 3/5

At the Plenary Meetings of Departments

807/30-58-8-9/43

new methods of detecting and reproducing signal: in the presence of interferences. Department of History: V. N. Lazarev, Corresponding Member, Academy of Sciences, USSR, spoke about the mosaics and frescoes of the St. Sophia Church in Kiyev and on the painting "The Yaroslav Panily". Department of Literature and Philology: Ya. Ye. El'sberg, Doctor of Philological Sciences, and Professor S. I. Oshegov, the Director of the Sektor sovremennogo literaturnogo yasyka i kul'tury rechi Instituta russkogo yasyka Akademii nauk SSSR (Branch for Modern Literary Language and Study of Languages of the Institute of Russian Language, AS USSR) spoke about the present reactionary theories and the revisionian in literature (El'sberg) and on some of the tendencies of these theories (Oshegov). El'sberg's ideas were backed by V. Y. Yermilov, Doctor of Philological Sciences. Oshegov's lecture evoked a brisk debate among the following persons: S. I. Kotkov and V. G. Orlova, Doctors of Philology, V. M. Sidorov and B. V. Gornung, Candidates of Philology, B. A. Serebrennikov, Corresponding Member, Academy of Sciences, USSR,

Card 4/5

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-2"

At the Plenary Meetings of Departments

507/30-58-8-9/43

V. V. Vinogradov, Member, Academy of Sciences, USSR.

Card 5/5

. AUTHORS: Knunyants, I. L., Gambaryan, M. P. 80T/62-58-10-10/25 TITLE: Determining the Strength of the Bonds of Radicals With Sulfur in Unsymmetrical Bulfides by Means of a Destructive Bromination Method (Opredeleniye prochnosti svyasi radikalov s seroy v nesimmetrichnykh sul'fidakh metodom destruktivnogo bromirovaniya) PERIODICAL: Isvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk, 1958, Mr 10, pp 1219-1227 (USSR) ABSTRACT: The reaction of the carbonyl compounds with mercaptans in the presence of anhydrous hydrogen chloride (Refs 2-5) is widely used in the synthesis of & -chlorosulfides. The reaction with other compounds can not be used for the synthesis. Sulfides that have hydrogen in the β -position to the alkthic group form, however, in the chlorination a mixture of products of the further chlorination of vinyl ethers; the latter form intermediately; and are difficult to separate. The bromination of the sulfides has remained almost uninvestigated. Only in 1956 it was shown that in the treatment of dimethyl sulfide by bromine a very stable bromosulfonium salt is formed which in Card 1/2 the course of long boiling in carbon tetrachloride decomposes

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Determining the Strength of the Bonds of Radicals
With Sulfur in Unsymmetrical Sulfides by Means of a
Destructive Bromination Method

into two directions (Ref 16). The authors of the present paper showed that the bromination of \$\beta\$ -alkthio carboxylic acid derivatives leads to the cleavage of the C-S bond, with a bromine derivative and disulfide being formed. The reaction of the destructive bromination can be taken as characteristic feature of the strength of the bond of radicals with sulfur in unsymmetrical sulfides. The binding strength of the investigated radicals increases according to the order:

\[C_{\text{H}_2} \cdot CHCH_2 \cdot COHHC_6 \text{H}_5 \cdot CHCH_2 \cdot COHHC_6 \text{H}_1 \cdot C_6 \text{H}_5 \cdot CH_2 \cdot CHCH_2 \cdot COHHC_6 \text{H}_1 \cdot C_6 \text{H}_5 \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_3 \), \$C = \(-\cdot CH_2 \cdot COHHC_6 \text{H}_1 \cdot - C_2 \text{H}_5 \cdot \text{There are 4 tables and 20 references, 2 of which are Soviet.} \]

ASSOCIATION:

Institut elementoorganicheskikh soysdineniy Akademii nauk SSSR (Institute of Elementary Organis Compounds, Academy of Sciences, USSR)

SUBMITTED: Card 2/2

March 4, 1957

40° 20° 与特别是中国国际的副科学的对照,但是**现在现在由部**的

5(3) AUTHORS:

Enunyante, I. L., Sterlin, R. N., Yatsenko, R. D., Pinkins, L. N.

807/62-58-11-11/26

TITLE:

Reactions of Fluoro Olefins (Reaktsii ftorolefinov) Communication VIII. Reactions of Perfluoro Vinyl Magnesium

Halides (Scobshoheniye 8. Reakteii

perftorvinilmagniygalogenidov)

PERIODICAL:

Isvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk, 1958, Hr 11, pp 1345-1347 (USSR)

ABSTRACT:

In the present paper the authors demonstrated that by the activation of magnesium with ethyl bromide and by carrying out the reaction in ester at -30 to -200 a practically quantitative consumption of magnesium can be achieved. By the decomposition of the reaction mass with diluted sulfurio acid 70 % of trifluoro ethylene could be separated. It was demonstrated that under the mentioned conditions perfluoro vinyl bromide and perfluoro vinyl chloride do not react with magnesium and that they are unchanged after the end of the reaction. An organomagnesium compound CF2-CFMgBr in a yield of up to 45 % could be formed from perfluoro vinyl bromide in tetrahydro furan. In this case it was not even necessary to

Card 1/3

APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-2"

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Reactions of Fluoro Olefins. Communication VIII. Reactions of Perfluoro Vinyl Magnesium Halides

507/62-58-11-11/26

activate magnesium with ethyl bromide. Apparently the assertion that an intensification of the basicity of the solvent favors the formation of R_pMgJ on the basis of its stabilization in the form of a complex of the



type, is justified. As the result of the processing of CF2 CFMgJ with solid carbon dioxide in ester solution at -40° and the subsequent decomposition of the reaction mass with 2N sulfuric acid solution perfluoro acrylic acid was obtained in a yield of 40 %. Henne (Ref 6) formerly obtained this acid by a complex and very slow method. The found method can be recommended without doubt for preparation. By processing the ester solution of the perfluoro acrylic

Card 2/3

Reactions of Fluoro Olefine. Communication VIII. Reactions of Perfluoro Vinyl Magnesium Halides

2 Kin A 2 Kin

50V/62-58-11-11/26

acid with a calculated amount of diamonethane the methyl ester of perfluoro acrylic acid was obtained. There are 8 references, 1 of which is Soviet.

SUBMITTED:

March 4, 1957

Card 3/3

(A) 1 大型(A) 1 (A) 2 (A)

5(3) AUTHORS:

Kil'disheva, O. V., Lin'kova, M. G., SOV/62-58-11-12/26 Savosina, V. M., Knunyants, I. L.

TITLE:

A. A. Disubstituted & Acylamino Carboxylic Acids (&, & -Disameshchennyye & -atsilaminokarbonovyye kisloty) Communication II. A New Method of Forming Oxazole-4-Carboxylic Acids (Soobshcheniye 2. Novyy sposob obrazovaniya oksazol-4-karbonovykh kislot)

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk, 1958, Nr 11, pp 1348-1353 (USSR)

ABSTRACT:

It has been communicated (Ref 1) that &, \$\beta\$-dihalogen-\$\times \text{-acylamino} propionic acids ensity react with water, alcohols, and amines and that they form &-substituted &\times \text{-acylamino}-\$\beta\$-halogen carboxylic acids (I). Further investigations have demonstrated that &, \$\beta\$-dihalogen-&-acylamino propionic acids easily react with mercaptans and according to the halogen (chlorine or bromine) mono- or dialkthic acids are obtained. In this paper a new reaction &-acylamino-\$\beta\$-halogen acrylic acids from &-acylamino-\$\beta\$-halogen acrylic acids is demonstrated. &-substituted &-acylamino-\$\beta\$-halogen propionic acids

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Communication II. A New Wethod of Forming Oxazole-4-Carboxylic Acids

307/62-58-11-12/26

transform into exampline carboxylic acids under the action of alkali. They form according to the conditions either acyloxy-pyroracemic acids or example carboxylic acids. The mechanism of formation of scyloxy pyroracemic acid from cc-substituted cc-acylamino- \(\beta\)-halogen carboxylic acids has been described already earlier (Ref 3). In this paper a mechanism of formation of example carboxylic acids from cc-acylamino- \(\beta\)-halogen acrylic acids was suggested. It was shown that the formation of example carboxylic acid from cc-substituted cc-acylamino- \(\beta\)-halogen propionic acids takes place over a stage of formation of 2-aryl (or alkaryl)-4-substituted example carboxylic acids without preceding transition into the corresponding cc-acylamino- \(\beta\)-halogen acrylic acids. There are 7 references, 3 of which are Soviet.

ASSOCIATION:

Institut elementoorganicheskikh soyedineniy Akademii nauk SSSR (Institute of Elemental organic Compounds of the Academy of Sciences, USSR)

Card 2/3

5(3) AUTHORS:

Kil'disheva, O. V., Shokina, V. V., Knunyants, I. L.

TITLE:

 α,β -Disubstituted α -Acylamino Carboxylic Acids $(\alpha,\beta$ -Dizameshchennyye- α -atsilaminokarbonovyye kisloty) Communication 3: α,α -Diacylamino- β -Halogen Propionic Acids (Soobshcheniye 3. α,α -Diatsilamino- β -galoidopropionovyye kisloty)

PERIODICAL:

Izvestiya Akademii nauk SSSR; Otdeleniye khimicheskikh nauk, 1958, Nr 12, pp 1461-1467 (USSR)

ABSTRACT:

Some time ago it was proved (Ref 1) that α, β -dihalogen- α -acylamino propionic acids (I) easily react with water, alcohols, amines and mercaptans, forming the corresponding α -substituted α -acylamino- β -halogen carboxylic acids (II). Unsuccessful attempts were made to obtain the α, α -diacylamino- β -halogen propionic acids (III), where X = NHCOR, by a reaction of α, β -dihalogen- α -acylamino propionic acids with the corresponding amides. Further investigations showed, however, that α, α -diacylamino- β -halogen propionic acids can easily be obtained by the condensation of halogen pyroracemic acids with the corresponding nitriles in the presence of concentrated sulfuric acid in much the same way as in the production of α, α -diacyl-

Card 1/2

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a, β -Disubstituted a-Aoylamino Carboxylic Acids. Communication 3: a, a-Discyl-amino- β -Halogen Propionic Acids

amino carboxylic acids (Ref 2). According to this method the α,α -diacylamino- β -halogen propionic acids mentioned in table 1 were obtained. The dehydration of α,α -diacylamino- β -halogen propionic acids on heating led to the saturated examples (VII) (Table 2). The example obtained were usually crystallized from accide anhydride as stable, colorless, crystalline compounds. On the action of methyl alcohol on 2-mothyl-4-acetylamino-4-chloro-methyl examples in the absence of moisture, the new α -amino- α -acetylamino- β -phloro propionic acid (VIII) with the melting point 1350 was easily formed. There are 2 tables and 7 references, 3 of which are Soviet.

ASSOCIATION:

Institut elementcorganicheakikh soyedineniy Akademii nauk SSSR (Institute of Elementorganic Compounds, Academy of Sciences,

SUBMITTED:

March 29, 1957

Card 2/2

AUTHOR:

Knunyants, I. L., Member, Academy of

807/29-58-10-10/28

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TITLE:

Scientists Great the Kemsenel !(Uchenyye privetstvuyut kemsemel!)Den't Get Clesed up in Yeur Special Field of Science (Ne samykaytes' v predelakh sveyey spetsial'nesti)

PERIODICAL:

Tekhnika meledezhi, 1958 Ar 1e, pp 12 - 12 (USSR)

ABSTRACT:

Among either things the auther quotes: Life means work and study. Indifference and passivity are the fate of the weak in mind. Indifference and passivity are no mode the fate of the eld they mean death. Learn, work and seek. Nothing is allowed to remain misunderstood and unsaid. From the organition of the less important the great and new is born. This is equal in life and science. Hencur knewledge and authority but do not bow to them. Think before you criticise. First of all you have to go to the bottom of a thing and then you are to defend it without the least fear and to express your epinion with courage! Be always honest! There is no greater sin than that of a biased idea, or to say semething one is not sure about only for reasons of preetige and influence. There

Card 1/2

Scientists Greet the Komsemel! Den't Get Clesed up in SOV/29-58-1e-1e/28

is no field of science which is not interesting. In every field of science great inventions are possible. In the mement a person starts to think indepedently his interest for science is roused. The individual fields of science are autonomous and closely linked. Do not restrict yourselves to the limits of your field of sciences! Do not close your minds to life: Be aware that philosophy is all-comprising. Without philosophy it welld not have been possible to discover the fundamental principles of life and development. Aquire the basic method of knewledge — materialistic dialectic — and you will always be successful in your work. There is 1 figure.

Card 2/2

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5(3) Auteors

Knunyants. I. L., Gambaryan, H. P., SOV/74-27-12-1/4 Rokhlin, Ye. M. (Moscow)

TITLE:

Carbenes (Karbeny) Compounds of Bivalent Carbon Occurring in Intermediary Form in Organic Reactions (Soyedineniya dvukhvalentnogo ugleroda, promeshutochno obrasuyushchiyesya v organicheskikh reaktsiyakh)

PERIODICAL:

Uspekhi khimii, 1958, Vol 27, Br 12, pp 1361 - 1436 (USSR)

ABSTRACT:

In this survey the authors made an attempt to deal thoroughly with the data known from publications on the intermediary formation of carbenes. In future the intermediary formation of carbenes may be expected to be demonstrated in the case of reactions of organic substances as well. The survey is concluded with the discussion of the carbene structure. At the moment it is not yet possible to say anything definite about the electron state of the carbenes - whether in singlet or triplet state. In the former case they can really be regarded as bases conjugate with carbon ions, in the latter case as radicals. The data in the publications are extremely contradictory. It frequently occurs that the individual authors draw different conclusions from one and the same

Card 1/4

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Carbones. Compounds of Bivalent Carbon Occurring as Intermediary Form in Organic Reactions

SOY/74-27-12-1/4

condition. Approximative quantum-mechanical computations lead, however, to the conclusion that the basic state of the most simple carbene - methylene - is a triplet state. The interest for carbenes was roused in connection with the work carried out with carbene dihalides. It was proved that in the case of an effect of bases on "haloforms" a separation of the proton takes place. The tribalogen methyl anion formed in this connection is decomposed into carbene dihalide and halogen anion. As a result of its electrophilic nature carbene dihalide enters a reaction with a number of nucleophilio reagents. In consequence of the reaction of carbene dihalides with olefins propane dihalides are formed. This new reaction has found a wide field of application and makes various cyclohexane derivatives accessible; among them also compounds with a condensed system containing a cyclopropane cycle. It is possible to explain the relative stability of carbones by means of the superposition of the following structures:

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and



of oxycarbenes may be explained by an unseparated pair of electrons in the "carbene"-carbon atom with the carbonyl unreliable in the synthesis of the explained by an unseparated pair of double binding. The carbone formation happens to be most APPROVED FOR RELEASE: 06/19/2000 CINTEDPROCESSING ON THE CARBON OF THE CARBON OF

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Carbenes. Compounds of Bivalent Carbon Occurring as Intermediary Form in Organic Reactions

807/74-27-12-1/4

derivatives. The possibility of and-separation of hydrogen halide is proved by the investigation of hydrogen halide separation of deutero halides of the type $\mathrm{RCD}_2\mathrm{CH}_2\mathrm{X}$ and $\mathrm{RCH}_2\mathrm{CD}_2\mathrm{X}$, even if the hydrogen atom is in a β -position. The d-separation of hydrogen halide is often accompanied by a process of regrouping which is in connection with the transformation of both hydrogen or deuterium and various groups connected with the β -hydrocarbon atom. Finally it may, however, be said that neither the geometric nor the electron structure of carbenes seems to be definitely investigated. There are 545 references, 72 of which are Soviet.

Card 4/4

20-119-1-22/52

AUTHORS: Golubeva, N. Ye., Kil'disheva, O. V., Knunyants, I. L.,

Member of Academy of Sciences

TITLE: Cancerolytic Peptides (Kantseroliticheskiye peptidy)

PERIODICAL: Doklady Akademii Nauk SSSR, 1958, Vol. 119, Nr 1,

pp. 83 - 86 (USSR)

ABSTRACT: Cancerolytic, sarcolysin-containing dipeptides (table 1) were produced by condensation of the M-formyl-derivative of

p-Mi-(A -ethyl chloride-)-amino-DL-phenylalanin (sarcolysin) with ethers of various amino acids. By the interaction of the

sarcolysin-ethyl-ether with p-di(\$-ethyl chloride)-amino-

phenyl-acetic-acid the ethyl ether of p-di-(8-ethyl chloride)-

aminophenacetyl-sarcolysin was obtained. Further p-di-(A-ethyl chloride)-aminophenacetyl- and y -[p-di-(A-ethyl

chloride)-amino J-phenylbutyryl-derivates of various amino

acids were produced whose general formula is given (table 2). By the interaction of p-di-(\beta-ethyl chloride)-aminophenyl-

Card 1/3 acetic-acid and \(\frac{1}{p-di-(\beta-ethyl chloride)-amino/phenylbutyric-

Card 1/5

20-119-1-22/52

Cancerolytic Peptides

acid with anilin in the presence of 1,3-dicyclo-hexyl-carbodiimide the corresponding anilides were obtained; it is true that in the case of the interaction of these acides with 2-methyl-5-ethoxymethylene-6-amino-pyrimidin or with p-di-(A-ethyl chloride)-aminophenyl-acetic-acid with cyclohexalamine only N-acyl-derivatives of 1,3-dicyclchexyl-urea were isolated. Preliminary samples on the cancerolytic action of the sarcolysin-containing peptides were performed in the Institute for Experimental Pathology and Cancer Therapy of the Academy of Medical Sciences of the USSR (Institut eksperimental'noy patologii i terapii raka Akademii meditsinskikh nauk SSSR). It became evident that the sarcolysin-peptides possess marked properties close to those of sarcolysin which act against tumors and which at the same time are not toxic and have a high selectivity of the action upon some tumors. Finally the general method of production of sarcolysin-containing peptides and that of the individual peptides is given in a kind of experimental part.

Card 2/3

20-119-1-22/52

Cancerolytic Peptides

There are 3 tables, and 1 reference, 1 of which is Soviet.

ASSOCIATION: Institut elementoorganicheskikh myedineniy Akademii nauk SSSR (Institute for Elementary Organic Compounds AS USSR)

SUBMITTED: December 10, 1957

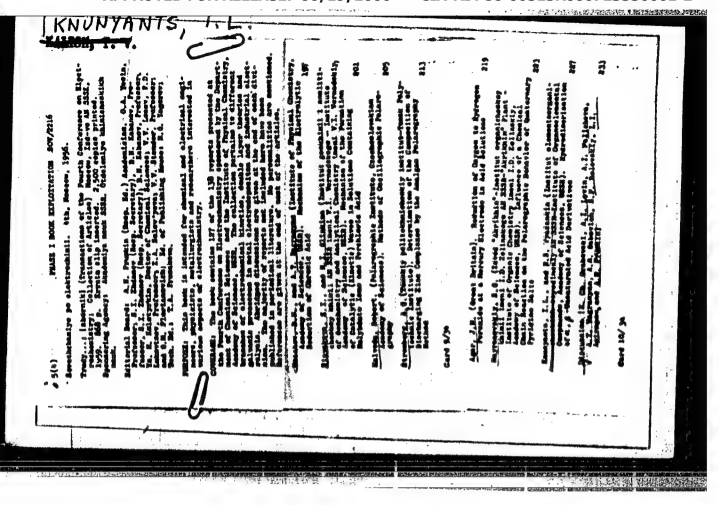
Card 3/3

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NHEMEYANOV, Aleksandr Mikolayevich, akademik; REUTOV, O.A., otv.red.town;
TOPCHIYEV, A.V., akademik, red.; EMUNYANTS, I.L., akademik, red.;
KABACHNIK, M.I., akademik, red.; FREYDMIER, R.EMT, red.; KAN, H.I.,
red.; LOEKUTOVA, I.P., red.isd-va; POLYAKOVA, T.V., tekim.red.

[Selected works in four volumes] Isbrannye trudy v chetyrekh tomakh. Moskva, Isd-vo Akad.nauk SSER. Vol.1. 1959. 712 p. (MIRA 12:12)

1. Ohleny-korrespondenty AN SESR (for Reutov, Freydline). (Chemistry)



APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-2"

KNUNYANIS, J.L.

PHASE I BOOK EXPLOITATION 80V/3494

- Reaktsii i metody issledovaniya organicheskikh soyedineniy, Kn. 8 (Beactions and Research Methods of Organic Compounds, Bk. 8) Moscow, Goskhimizdat, 1959. 446 p. Errata slip inserted. 4,200 copies printed.
- Eds (Title page): V.M. Sodionov, Academician (Deceased), B.A. Kazanskiy, Academician, I.L. Knunyants, Academician, M.M. Shenyakin, Academician, and H.M. Mel'nikov, Professor; Ed. (Inside book): V.P. Yevdakov; Tech. Ed.: V.F. Zazul'skaya.
- PURPOSE: This book is intended for laboratory personnel at industrial plants, for instructors and students at higher educational establishments, and particularly for scientists and researchers working at the numerous research institutes in the Soviet Union.
- COVERAGE: This is the eighth volume in a series "Reactions and Research Methods of Organic Compounds." This series does not duplicate the one published in English under the title "Organic Reactions" and appearing in Bussian translation; however, some material, of particular interest, is included in this publication. The present series is primarily devoted to reviewing the works of Boviet chemists. The eighth volume of this series deals with thiocyanation

Card 1/5

Reactions and Research (Cont.)

804/3494

reactions of organic compounds and methods of studying them. It presents data on analytical methods using thiocyanates for the study of fats, mineral oils, and volatile oils. It discusses the use of thiocyanates for photographic emulsions, acceleration of rubber vulcanization, stabilization of lubricating oils, purification of tars, abstement of corrosion and purification of metals, production of mustard oil, and synthesis of sulfur-bearing compounds. It also discusses the use of thiocyanates as solvents for acrylonitrile polymers, as intermediate products in the synthesis of dyes, as antiseptics, bactericides, medicines, insecticides, and fungicides. The book contains 327 pages of tables listing 1449 initial organic compounds subjected to thiogyanation. The tables give formulas of the initial compounds, the names and structural formulas of the compounds, the reaction conditions, the reaction products and their yield percent as compared with the theoretical yield, as well as references to the literature on which the data are based. There are 797 references: 376 English, 228 German, 74 Soviet, 47 French, 17 Italian, 25 Japanese, 7 Polish, 7 Scandinavian, 3 Belgian, 8 Swiss, 1 Dutch, and & Bungarian.

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5(3) AUTHORS:

Sterlin, R. W., Sidorov, V. A., Knunyanta, I. L.

TITLE:

Reactions of Fluoro Olefins (Reaktsii ftorolefinov)
Communication IX. Action of Anhydrous Aluminum Trichloride
on Fluoro Olefins (Soobshcheniye 9. Deystviye besvodnogo
trekhkhloristogo alyuminiya na ftorolefiny)

PERIODICAL:

Izvestiya Akademii nauk SSSR. Otdeleniye khimicheskikh nauk, 1959, Nr 1, pp 62 - 64 (USSR)

ABSTRACT:

In the present paper the authors investigated the effect exercised by anhydrous AlCl₃ on fluorinated elefins and especially on perfluoro propylene and 2-hydroperfluoro propylene. As a result of the interaction of CP₃— CH — CP₂ with AlCl₃ pentachloro propene CCl₃— CH — CCl₂ is synthesized as the only reaction product in a CH₃COCl solution under pressure (yield 65.5%). Similar results were obtained in the cold, at atmospheric pressure and in the substitution of chloroform for chloro acetyl. The complete exchange of fluorine atoms for chlorine in fluorinated elefins takes

Card 1/3

Reactions of Fluoro Olefins. Communication IX. Action of Anhydrous Aluminum Trichloride on Fluoro Olefins

807/62-59-1-9/38

place under extremely soft conditions. This exchange apparently is a result of the σ , a conjugation in the 2-hydroperfluoro propylene molecule

which determines the mobility of fluorine atoms of the GP, group and the levity of the silyl regrouping. In the reaction of AlCl, with perfluoro propylene, which was carried out under equal conditions as in the case of 2-hydroperfluoro propylene, the only reaction product obtained was a compound with a C₃FCl₃ composition. Its structure may be expressed by one of the following formulae: CFCl₂— CCl — CCl₂, CCl₃— CF — CCl₂ (Ref. 3). The compounds obtained were exidised in order to determine their structure. Trichloroacetic gold was synthesized as a result of the oxidation.

Card 2/3

"APPROVED FOR RELEASE: 06/19/2000 CIA-RDP86-00513R000723330001-2

Reactions of Fluoro Olefins. Communication II. Action of Anhydrous Aluminum Trichloride on Pluoro Olefine

507/62-59-1-9/38

This may be taken as a proof that 1,1,1,3,3-pentachloro-2-fluoropropylene-3 was obtained as a result of the exchange.
There are 3 references, 1 of which is Soviet.

SUBMITTED:

April 17, 1957

Card 3/3

5 (0) AUTHORS: Knunyants, I. L., Topchiyer, A. T. 507/62-59-8-2/42 TITLE: On the Occasion of A. H. Nesmejanov's 60th Birthday PERIODICAL: Izvestiya Akademii nank SSSR. Ctdeleniye khimicheakikh nank, 1959, Nr 8, pp 1357-1361 (USSR) ABSTRACT: The editors of the present journal and the Department of Chemical Sciences of the Academy of Sciences, USSR (Otdeleniye khimicheskikh nauk Akademii nauk SSSR) congratulate A. N. Nesmeyanov on his 60th birthday, which he celebrated on September 9, 1959. This great Soviet scientist is then celebrated for his scientific work. Wesneyanov had graduated from Moskovskiy gosudarstvemmy universitet (Moscow State University) at the early age of 23 and had then worked as Assistant, Docent, and holder of the Chair of Organic Chemistry, and had finally become the Rector of this university, the oldest of the country. He was among the founders of the new university on the Lambachye gory. The following scientific studies are mentioned: The method of synthesising organometallia compounds (1929) which now bears the name of Mesmeyanov resotion. In his research work he was mainly interested in the Card 1/3 problems of the borderline between and transition of inorganic

On the Occasion of A. W. Mesmeyanov's 60th Birthday

507/62-59-8-2/42

and organic ormpounds. He showed that the metal-carbon bond is unstable if an a- or f-valence level is not utilized. He found in the investigation of the orientation of substitution in a ferrocene moleus that the conjugated bond is transferred in the molecule by the iron. He also succeeded in determining some rules governing tautomeries and twofold reaction possibility of organic substances while studying organometallic compounds. In his work concerning the stersochemistry of the ois- and trans isomer of arsenio, antimony, and other metalchloro-\$-vinyl derivatives he obtained a considerable understanding of the relationship between the configuration of a compound and its physiological effect. Thanks to this interest in the chamistry of elements with variable or uncommon valencies and owing to the investigation of their carbonyl ocupounds he succeeded in developing methods for the preparation of these compounds in their absolutely pure form, Among his research work in the field of telomerisation of various bifunctional compounds the synthesis of the «-amino-enanthic acids deserves special mention, by whose polymerization he arrived at a new kind of polyamide fibers, the "enant fibers". On the basis of these investigations a method for the industrial

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On the Occasion of A. W. Nemmeyanov's 60th Birthday SOV/62-59-8-2/42

production of such fibers was developed at the Institut elementoorganicheskikh soyedineniy AN SSSR (Institute of Elemental-organic Compounds of the Academy of Sciences, USSR) in cooperation with GIAP. Investigations of the so-called inner transformation of molecules (radical isomerisation) let him discover new laws of nature. Moreover, Nesseyanov devoted his attention to a number of problems of scientific organisation and actively promoted scientific cooperation. There is 1 figure.

Card 3/3

5.3600 77290 SOV/63-4-6-24/37 AUTHORS: Sterlin, R. N., Pinkina, Yatsenko, R. D., Knunyants, I. L. TITLE: Brief Communications. Perfluorovinyl Derivatives of As PERIODICAL: Khimicheskaya nauka i promyshelnnost', 1959, Vol 4, Nr 6, pp 800-801 (USSR) ABSTRACT: Tertiary perfluorovinyl derivatives of As and Sb were obtained by the reaction of perfluorovinylmagnesium iodide with AsCl3 and SbCl3 in ether solution. Primary and secondary perfluorovinylarsine and corresponding stibine were not obtained. Perfluorovinyl dichloroarsine was obtained by the reaction of 10-alky1-5-10--dihydrophenarsazine and liquid HCl. The corresponding perfluorovinyl derivative of dihydrophenarsazine was obtained from perfluorovinylmagnesium iodide and adamsite. Perfluorovinyldichloroarsine (a new product) was obtained by decomposition of 10-perfluoroviny1-5,10-dihydrophenarsazine with liquid HCl. The obtained substances have the following properties: tri-(trifluorovinyl)-Card 1/2

Brief Communications. Perfluorovinyl Derivatives of As and Sb

77290 SOV/63-4-6-24/37

-arsine, bp 58°/95 mm, 50°/70 mm, and 110-111°/746 mm, 18 1.3938, d₄ 18 1.8400, in 40% yield. Tri-(trifluoro-viny1)-stibine, bp 75-75.5°/74 mm, n_D 24 1.4190, d₄ 2.06, in 41% yield. 10-Trifluoroviny1-5,10-di-hydrophenarsazine, mp 122° (alcohol) in 75% yield. Trifluoroviny1 dichloroarsine, bp 115°, n_D 20 1.4820, d₄ 1.9800, in 92.5% yield. There are 2 references, 1 Soviet, 1 French.

SUBMITTED:

June 1, 1959

Card 2/2

5.3600

77292

SOV/63-4-6-26/37

AUTHORS:

Knunyants, I. L., Rokhlin, E. M., Gambaryan, N. P., Cheburkov, Yu. A., Ch'en sh'ing-yun

TITLE:

Brief Communications. Fluorinated Ketones. Bis-

(trifluoromethyl)-glycolia Acid

PERIODICAL:

Khimicheskaya nauka 1 promyshlennost', 1959, Vol 4, Nr 6, pp 802-804 (USSR)

ABSTRACT:

Nitrile of bis-(trifluoromethyl)-glycolic acid (I) was synthesized by the reaction of hexachloroacetone with HCN in the presence of a catalytic amount of

piperidine.

(CF3)2CO + HCN piperidine (CF3)2CCOH (I)

(I) can be distilled at atmospheric pressure without decomposition but, in the presence of piperidine, (I) is decomposed to HCN and hexafluoroscetone. (I) is partially hydrolyzed in the presence of water at room temperature, forming hexafluoracetone hydrate and HCN.

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